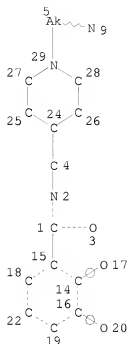


L6 HAS NO ANSWERS
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NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
DEFAULT ELEVEL IS LIMITED

GRAPH ATTRIBUTES:
RING(S) ARE ISOLATED OR EMBEDDED
NUMBER OF NODES IS 20

STEREO ATTRIBUTES: NONE

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100.0% PROCESSED 1283 ITERATIONS 194 ANSWERS
SEARCH TIME: 00.00.01

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COST IN U.S. DOLLARS SINCE FILE TOTAL
ENTRY SESSION
FULL ESTIMATED COST 360.86 361.07

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FILE COVERS 1907 - 28 Oct 2008 VOL 149 ISS 18
FILE LAST UPDATED: 27 Oct 2008 (20081027/ED)

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L8 5 L7

=> d bib abs 1-5

L8 ANSWER 1 OF 5 CAPLUS COPYRIGHT 2008 ACS on STN

AN 2005:34761 CAPLUS

DN 142:113915

TI Preparation of heterocyclic substituted 4-(aminomethyl)piperidine benzamides as 5-HT4 antagonists

IN Bosmans, Jean-Paul Rene Marie Andre; Gijssen, Henricus Jacobus Maria; Mevellec, Laurence Anne

PA Janssen Pharmaceutica N. V., Belg.

SO PCT Int. Appl., 46 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 2005003124	A1	20050113	WO 2004-EP6273	20040610
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	RW:	BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG			
	AU 2004254190	A1	20050113	AU 2004-254190	20040610
	CA 2528642	A1	20050113	CA 2004-2528642	20040610
	EP 1638961	A1	20060329	EP 2004-739776	20040610
	R:	AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, PL, SK, HR			
	JP 2006527714	T	20061207	JP 2006-515879	20040610
	US 20070197600	A1	20070823	US 2005-560486	20051212
PRAI	WO 2003-EP50240	A	20030619		
	WO 2004-EP6273	W	20040610		

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB The title compds. I [R1R2 = O(CH2)nO, n = 1-4; R1R2 = O(CH2)m, m = 2-5; R3 = H, halide, C1-C6-alkyl, C1-C6-alkoxy; R4 = H, halide, C1-C6-alkyl, C1-C6-alkoxy, cyanoalkyl, NH2, mono or di(C1-C6-alkyl)amino; R5 = H, C1-C6-alkyl, OR5 = 3- or 4-position; L = C1-C12-alkanediyl-R6, C1-C12-alkanediyl-X-R7; R6, R7 = heterocycle or heterocycle substituted with halide, OH, C1-C6-alkyl; heterocycle = morpholine, tetrazole, pyrazole, isoxazole, isothiazole, oxazolyl, thiazole, pyran, 2,4-dioximidazolidine] were prepared and tested as 5-HT4 antagonists. For example, reacting (chloropropyl)trityltetrazole II with trans-hydroxypiperidine derivative III gave alkylated piperidine IV (R = CPh3) which was deprotected to give IV (R = H).

RE.CNT 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 2 OF 5 CAPLUS COPYRIGHT 2008 ACS on STN

AN 2005:34760 CAPLUS

DN 142:134469

TI A preparation of 5HT4-antagonistic N-(piperidin-4-ylmethyl)-benzamide derivatives

IN Bosmans, Jean-Paul Rene Marie Andre; Gijssen, Henricus Jacobus Maria; Mevellec, Laurence Anne

PA Janssen Pharmaceutica N. V., Belg.

SO PCT Int. Appl., 63 pp.

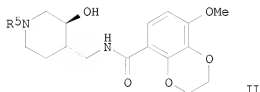
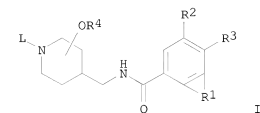
CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 2005003122	A1	20050113	WO 2004-EP6278	20040610
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	RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
	AU 2004254193	A1	20050113	AU 2004-254193	20040610
	CA 2528656	A1	20050113	CA 2004-2528656	20040610
	EP 1638959	A1	20060329	EP 2004-736513	20040610
	EP 1638959	B1	20070919		
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	JP 2006527715	T	20061207	JP 2006-515883	20040610
	AT 373651	T	20071015	AT 2004-736513	20040610
	ES 2293262	T3	20080316	ES 2004-736513	20040610
	US 20070032486	A1	20070208	US 2005-560485	20051212
PRAI	WO 2003-EP50237	A	20030619		
	WO 2004-EP6278	W	20040610		



AB The invention relates to a preparation of 5HT4-antagonistic N-(piperidin-4-ylmethyl)-benzamide derivs. of formula I [wherein: R1 is a bivalent (un)substituted radical of formula O(CH2)1-4O or O(CH2)2-5; R2 is H or halogen; R3 is alkyl, alkoxy, or halogen; R4 is H or alkyl, and OR4 is situated at the 3- or 4-position of the piperidine ring; L is H, alkyl-(H/OH/CN/cycloalkyl), alkyl-(O/S/SO2)-(cyclo)alkyl, or alkyl-C(O)-alkyl, etc.]. For instance, N-(piperidin-4-ylmethyl)-benzamide derivative II (R5 = H; pIC50 = 6.97) was prepared via decarboxylation of II (R5 = t-BuO2C).

RE.CNT 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 3 OF 5 CAPLUS COPYRIGHT 2008 ACS on SIN
AN 2005:14394 CAPLUS
DN 142:114101

TI Preparation of N-(piperidinylmethyl) benzamide derivatives as 5HT4-antagonists

IN Bosmans, Jean-Paul Rene Marie Andre; Gijssen, Henricus Jacobus Maria; Mevellec, Laurence Anne

PA Janssen Pharmaceutica N.V., Belg.

SO PCT Int. Appl., 98 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 200500838	A1	20050106	WO 2004-EP6285	20040610
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RW:	BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, BG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE,			

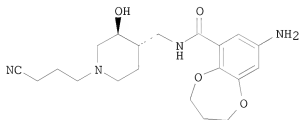
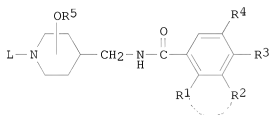
SN, TD, TG

AU 2004251825	A1	20050106	AU 2004-251825	20040610
CA 2528590	A1	20050106	CA 2004-2528590	20040610
EP 1641784	A1	20060405	EP 2004-739785	20040610
EP 1641784	B1	20070613		

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CN 1809558	A	20060726	CN 2004-80017134	20040610
BR 2004011603	A	20060808	BR 2004-11603	20040610
JP 2006527716	T	20061207	JP 2006-515886	20040610
AT 364602	T	20070715	AT 2004-739785	20040610
ES 2288258	T3	20080101	ES 2004-739785	20040610
NZ 544592	A	20080530	NZ 2004-544592	20040610
US 20060281753	A1	20061214	US 2005-560479	20051212
MX 2005PA13772	A	20060308	MX 2005-PA13772	20051215
IN 2005DN06108	A	20071214	IN 2005-DN6108	20051228
PRAI WO 2003-EP50236	A	20030619		
WO 2004-EP6285	W	20040610		

OS CASREACT 142:114101; MARPAT 142:114101
GI



AB Title compds. represented by the formula I [wherein R1R2 = OCH2O, O(CH2)nOm, O(CH2)5; n = 2-4; m = 0 or 1; R3 = H, halo or alkyl; R4 = (cyano)alkyl, alkoxy(alkyl), cyano, (alkyl)amino; R5 = H or alkyl; L = H, alkyl(cyano), alkoxyalkyl, alkylcarbonyl, etc.; stereochem. isomers thereof, an N-oxides thereof, and pharmaceutically acceptable acid or base addition salts thereof] were prepared as 5HT4-antagonists. For example, II was given in a multi-step synthesis starting from Me

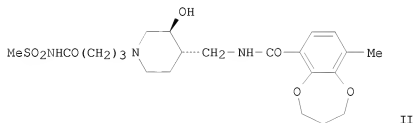
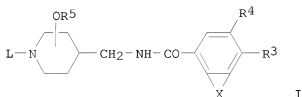
5-nitro-2,3-dihydroxybenzoate. I were tested for 5HT4 antagonistic activity with pIC50 values of around 6-9, and showed metabolic stability as well. Thus, I and their pharmaceutical compns. are useful as a medicine of 5HT4-antagonists.

RE.CNT 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 4 OF 5 CAPLUS COPYRIGHT 2008 ACS on STN
AN 2005:14393 CAPLUS
DN 142:113910

TI Preparation of aminosulfonyl substituted 4-(aminomethyl)-piperidine
 benzamides as 5HT4-antagonists
 IN Bosmans, Jean-Paul Rene Marie Andre; Gijssen, Henricus Jacobus Maria;
 Mevellec, Laurence Anne
 PA Janssen Pharmaceutica N.V., Belg.
 SO PCT Int. Appl., 53 pp.
 CODEN: PIXXD2
 DT Patent
 LA English
 FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 2005000837	A1	20050106	WO 2004-EP6280	20040610
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	RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
	AU 2004251823	A1	20050106	AU 2004-251823	20040610
	CA 2526079	A1	20050106	CA 2004-2526079	20040610
	EP 1641783	A1	20060405	EP 2004-739781	20040610
	R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, PL, SK, HR				
	JP 2007526875	T	20070920	JP 2006-515884	20040610
	US 20060142339	A1	20060629	US 2005-560300	20051212
PRAI	EP 2003-50238	A	20030619		
	WO 2003-EP50238	A	20030619		
	WO 2004-EP6280	W	20040610		
OS	MARPAT 142:113910				
GI					



AB Novel compds. of formula I [X = O(CH2)nO, O(CH2)n; n = 1-5; R3 = H, halo, alkyl, alkoxy; R4 = H, halo, alkyl, alkoxy, cyanoalkyl, CN, (substituted) amino; R5 = H, alkyl; L = (substituted) aminosulfonylalkyl, alkylsulfonylaminocarbonylalkyl, etc.] are prepared which have

5HT4-antagonistic properties. The invention further relates to methods for preparing such compds., pharmaceutical compns. comprising said compds. as well as the use as a medicine of said compds. Thus, II was prepared, and had 5HT4 antagonism activity with pIC50 of 7.92, and was 5% metabolized after 60 min in liver tissue.

RE.CNT 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 5 OF 5 CAPLUS COPYRIGHT 2008 ACS on STN

AN 2000:441788 CAPLUS

DN 133:74035

TI Preparation of 4-(aminomethyl)piperidinebenzamides as gastrointestinal agents.

IN Bosmans, Jean-Paul Rene Marie Andre; Meulemans, Ann Louise Gabrielle; De Cleyn, Michel Anna Jozef; Gijzen, Henricus Jacobus Maria

PA Janssen Pharmaceutica N.V., Belg.

SO PCT Int. Appl., 58 pp.

CODEN: PIXXD2

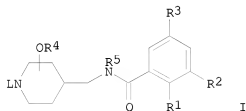
DT Patent

LA English

FAN.CNT 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2000037461	A1	20000629	WO 1999-EP10064	19991214
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TW 570920	B	20040111	TW 1999-88121125	19991203
CA 2355857	A1	20000629	CA 1999-2355857	19991214
BR 9916491	A	20010904	BR 1999-16491	19991214
EP 1140915	A1	20011010	EP 1999-967956	19991214
EP 1140915	B1	20050615		
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HU 2001004838	A2	20020729	HU 2001-4838	19991214
HU 2001004838	A3	20030528		
EE 200100335	A	20020815	EE 2001-335	19991214
JP 2002533337	T	20021008	JP 2000-589532	19991214
NZ 512871	A	20021126	NZ 1999-512871	19991214
AU 770397	B2	20040219	AU 2000-24328	19991214
IL 143858	A	20050320	IL 1999-143858	19991214
AT 297917	T	20050715	AT 1999-967956	19991214
PT 1140915	T	20051130	PT 1999-967956	19991214
ES 2245131	T3	20051216	ES 1999-967956	19991214
SK 285829	B6	20070906	SK 2001-859	19991214
PL 197409	B1	20080331	PL 1999-348417	19991214
IN 2001MN00442	A	20050304	IN 2001-MN442	20010423
BG 105571	A	20020131	BG 2001-105571	20010607
BG 64953	B1	20061031		
NO 2001002858	A	20010608	NO 2001-2858	20010608
NO 321324	B1	20060424		
US 6544997	B1	20030408	US 2001-857905	20010608
HR 2001000445	A1	20020630	HR 2001-445	20010614
MX 2001PA06409	A	20010910	MX 2001-PA6409	20010621
ZA 2001005135	A	20020621	ZA 2001-5135	20010621

HK 1039114	A1	20050819	HK 2002-100161	20020129
US 20030181456	A1	20030925	US 2003-353307	20030129
US 7205410	B2	20070417		
PRAI EP 1998-204411	A	19981222		
WO 1999-EP10064	W	19991214		
US 2001-857905	A3	20010608		
OS MARPAT 133:74035				
GI				



AB Title compds. [I; R1R2 = (substituted) OCH2O, OCH2CH2, OCH2CH2O, etc.; R3 = H, halo; R4, R5 = H, alkyl; L = cycloalkyl, oxocycloalkyl, alkenyl, etc.], were prepared. Thus, trans-N-[1-(3-aminopropyl)-3-hydroxy-4-piperidinylmethyl-7-chloro-2,3-dihydro-1,4-benzodioxin-5-carboxamide (preparation given). Was stirred with 2-chloro-3-methylpyrazine and CaO at 120° to give 16% trans-7-chloro-2,3-dihydro-N-[3-hydroxy-1-[3-[(3-methyl-2-pyrazinyl)amino]propyl]-4-piperidinylmethyl]-1,4-benzodioxin-5-carboxamide. This antagonized 5HT4 in rat esophageal tunica muscularis mucosae with pA2 = 10.55.

RE.CNT 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT